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DOCKET NO.: CACO-0067 (P21303US) PATENT APPLICATION

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IN THE CLAIMS:

Please amend claims 1 - 19 and add new claims 20 and 21 to appear as follows.

1 (Amended). A method of selecting a nucleic acid encoding an enzyme that is capable of converting a prodrug to its active drug form comprising the steps of:

- a) contacting a population of bacteria transformed with a bacteriophage library with a prodrug in a medium, wherein:
 - i) the transformed bacteria are in the lysogenic state, and
- ii) when converted to its active drug form, the prodrug causes activation of the proteolytic activity of bacterial RecA and lysis of the bacteria;
- b) separating bacteriophage particles released by lysis of the bacteria from said medium; and
- c) analysing the genotype of said released bacteriophage particles for a nucleic acid encoding the enzyme.
- 2 (Amended). A method of selecting a nucleic acid encoding an enzyme capable of converting a prodrug to its active drug form comprising the steps of:
- a) introducing a library of genes into bacteriophage to form a bacteriophage library;
 - b) infecting a population of bacteria with said bacteriophage library;
- c) selecting said infected bacteria for bacteria in which the lysogenic state has been established;
 - d) contacting said bacteria with said prodrug in a medium;
- e) separating from said medium bacteriophage particles released by lysis of host bacteria; and

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- f) analysing the genotype of said released bacteriophage for the nucleic acid encoding the enzyme;
- g) wherein said prodrug causes activation of the proteolytic activity of bacterial RecA when converted to its active drug form.
- 3 (Amended). The method of claim 1 or claim 2, wherein the steps are repeated in at least one cycle.
- 4 (Amended). The method of claim 1 or 2, wherein the genotype of said released bacteriophage particles is analysed by DNA sequencing.
- 5 (Amended). The method of claim 1 or 2, wherein said bacteriophage carry a gene encoding antibiotic resistance or other selectable marker.
- 6 (Amended). The method of claim 1 or 2, wherein said enzyme is selected from the group consisting of nitroreductase, flavin reductase, DT-diaphorase, thymidine kinase, cytosine deaminase, and purine nucleoside phosphorylase.
- 7 (Amended). The method of claim 1 or 2, wherein said prodrug is selected from the group consisting of CB1954, SN 23862, 2-[N,N-bis(2-iodoethyl)amino]-3,5-dinitrobenzamide, 5-fluorocytosine, acyclovir, ganciclovir, and 6-methyl-9-(2-deoxy-β-D-erythro-pentofuranosyl) purine.
- 8 (Amended). The method of claim 1 or 2, wherein said bacteriophage is the bacteriophage lambda or a lambda derivative.
- 9 (Amended). The method of claim 2, wherein said gene library comprises genes encoding variants of a single enzyme.

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- 10 (Amended). The method of claim 9, wherein said variants comprise amino acid deletions and/or insertions and/or substitutions from the wild type enzyme.
- 11 (Amended). The method of claim 9, wherein said genes encoding said variants are generated by DNA shuffling, random mutagenesis, or PCR shuffling.
- 12 (Amended). The method of claim 1 or 2, wherein said activity of said bacterial RecA protein is caused by the generation of single-stranded DNA in the bacterium.
- 13 (Amended). The method of claim 12, wherein said single-stranded DNA is generated as a consequence of the enzymatic conversion of the prodrug to its active drug form.
- 14 (Amended). The method of claim 12, wherein said single-stranded DNA is generated as a result of a break in one or both strands of the DNA, a cytotoxic lesion, a DNA crosslink or a monovalent DNA adduct, or by inhibition of the progress of DNA replication [by any other means].
- 15 (Amended). The method of claim 1 or 2, wherein said enzyme comprises nitroreductase and said prodrug comprises CB1954.
- 16 (Amended). The method of claim 1 or 2, wherein said bacteriophage is λJG3J1.
- 17 (Amended). The method of claim 1 or 2, wherein said bacteria are E. coli strain C600Hfl.
- 18 (Amended). A nucleic acid molecule encoding a catalytic enzyme or enzyme fragment isolated according to the method of claim 20 or 21.

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19 (Amended).

A catalytic enzyme or enzyme fragment encoded by the nucleic acid molecule

of claim 18.

20 (New). A method of cloning a nucleic acid encoding a catalytic enzyme or enzyme fragment, said catalytic enzyme or enzyme fragment being capable of converting a prodrug to its active drug form, comprising the steps of:

- a) contacting a population of bacteria transformed with a bacteriophage library with a prodrug in a medium, wherein
 - i) the transformed bacteria are in the lysogenic state, and
- ii) when converted to its active drug form, the prodrug causes activation of the proteolytic activity of bacterial RecA and lysis of the bacteria;
- b) separating bacteriophage particles released by lysis of the bacteria from said medium;
- c) analyzing the genotype of said released bacteriophage particles for a nucleic acid encoding the enzyme, or functional fragment thereof; and
- d) cloning the nucleic acid of the released bacteriophage particles that encode the enzyme or enzyme fragment.
- 21 (New). A method of cloning a nucleic acid encoding a catalytic enzyme or enzyme fragment, said catalytic enzyme or enzyme fragment being capable of converting a prodrug to its active drug form, comprising the steps of:
- a) introducing a library of genes into bacteriophage to form a bacteriophage library;
 - b) infecting a population of bacteria with said bacteriophage library;
- c) selecting said infected bacteria for bacteria in which the lysogenic state has been established;